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The temporal coordination of anaphase, cytokinesis and mitotic exit is essential for the production of viable daughter cells, and mutations that affect the proper timing of these events. result in genomic instability, a hallmark of cancer. In yeast, a signaling pathway has been that identified, called the Mitotic Exit Network, which coordinates mitotic exit and cytokinesis with the end of anaphase. The identification and characterization of such a pathway in human cells is necessary to further our understanding of how normal cell division is regulated and will highlight possible mechanisms of genomic instability in tumor cells. In order to discover those genes that are involved specifically in animal cell division and are not conserved in yeast, we are taking advantage of the nematode, C. elegans. Using the yeast two-hybrid system, we are currently mapping the physical interactions of many C. elegans genes believed to be potential mitotic regulators. The goal is to generate a large protein interaction map, which in combination with phenotypic data will help to elucidate the biochemical pathways involved in cooridinating mitotic events in animal cells. In work completed up to now, we have generated a preliminary mitotic protein-interaction map, which still requires stringent refining and confirmation based on additional protein interaction assays.

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· Introduction

The temporal coordination of anaphase, cytokinesis and mitotic exit is essential for the production of viable daughter cells, and mutations that affect the proper timing of these events result in genomic instability, a hallmark of cancer. In veast, a signaling pathway has been identified, called the Mitotic Exit Network, which coordinates mitotic exit and cytokinesis with the end of anaphase. Homologues of three of these signaling components have been identified in humans suggesting that human cells regulate mitosis in a similar fashion; however, a clear mitotic exit network has yet to be revealed. The identification and characterization of such a pathway in human cells will further our understanding of how normal cell division is regulated and will highlight possible mechanisms of genomic instability in tumor cells. In order to discover those genes that are involved specifically in animal cell division and are not conserved in yeast, we are taking advantage of the nematode, C. elegans. C. elegans is a multi-cellular complex metazoan whose genes are more homologous to humans than are those of yeast. This system will allow for the rapid functional analysis of large numbers of candidate genes that can then be used to ascertain their human counterparts by sequence comparison. Using the yeast two-hybrid system we have built a protein interaction map for hundreds of candidate C. elegans genes that are potentially involved in mitotic temporal control, based on phenotypic data as well as homology to yeast genes of known function. Once complete, these interaction data in combination with phenotypic data will help to elucidate specific biochemical pathways involved in late mitotic events. By recognizing the components of these putative pathways novel targets for anti-cancer therapies may be discovered. This report summarizes work done over the past twelve months, which culminated in the drawing of a preliminary mitotic protein interaction map for C. elegans.

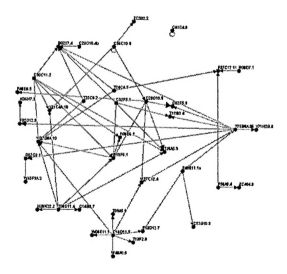
Body

- 1. The generation of constructs to be used in the yeast two-hybrid assay: As outlined in my proposal, I selected 324 *C. elegans* genes that were likely to be involved in mitotic regulation, based on RNAi phenotypes and homology to yeast genes of known function. I received clones for each gene from Marc Vidal, a collaborator at Harvard who has cloned the majority of *C.elegans* cDNAs into plasmids to create the worm "ORFeome" (ref 1). After receiving these clones from Dr. Vidal they needed to be subcloned into vectors for use in the yeast two-hybrid assay. The ORFeome was created using Invitrogen's Gateway vector system, which allows inserts to be transferred from one vector to another by site-directed recombination. By eliminating the need to subclone by the conventional restriction digest/ligation method, this system greatly reduced the amount of effort needed to transfer all 324 clones into both bait and prey vectors. Using a high-throughput technique I was able to successfully generate both bait and prey constructs for all 324 genes of interest. After the prey plasmids were constructed they were combined to create a normalized mitosis-specific library for two-hybrid screening. The bait plasmids were individually transformed into an appropriate yeast strain.
- 2. The generation of yeast strains to be used in the yeast two-hybrid assay: After the appropriate bait plasmids were obtained for each of the 324 genes, they were individually transformed into the yeast strain MaV203, and later into strain PJ69A, which contain the appropriate genetic requirements for yeast two-hybrid experiments. MaV203 has reporter genes HIS3, URA3, and LacZ, while PJ69A uses HIS3, ADE2, and LacZ reporters. Most derivative strains were made using a high-throughput yeast transformation protocol, described in reference 1. None of the bait constructs proved to be lethal to the yeast strains, and only two baits were able to auto-activate the reporter genes.
- 3. Screening yeast two-hybrid libraries: The first set of experiments were done by screening complete yeast two-hybrid cDNA libraries with baits of particular interest, such as the worm homologues of yeast Mitotic Exit Network components, MOB1 and CDC5. The libraries used were gifts from Dr. Vidal and included an ORFeome prey library (a collection of all plasmids currently in the ORFeome) and the cDNA library that the ORFeome was originally cloned from. We had no success at screening these libraries using the MaV203 derived strains due to extremely high background with the HIS3 reporter. We eventually switched to the PJ69A derived strains, which allowed us to select for two-hybrid interactions on medium lacking adenine, a much stronger selection than HIS3. Using this strain background a large number of two-hybrid interactions were obtained for each of the baits; however, after sequencing the prey plasmids we realized that there was a subset of preys that were hit in every screen, regardless of which bait was used. One common feature of

these non-specific prey plasmids was that they all encoded for proteins that contained cysteine-rich domains, or were enzymes involved in the modification of disulfide bridges. Because I was unable to reduce the level of this background without hampering my ability to detect real interactions, I decided not to continue with the deep two-hybrid screens and instead focused on screening the mitotic library that I had made because this library did not contain any of those unwanted prey plasmids.

- 4. Yeast two-hybrid matrix experiment: The original idea for the two-hybrid matrix was to test in pair-wise fashion the interaction between every bait construct and every prey construct in my candidate gene collection. This amounts to (324 x 324) 104.976 individual two-hybrid experiments. To make this experiment feasible we were planning to create two sets of yeast strains, one set expressing each of the bait plasmids (MaV203 derived) and the other set expressing the prey plasmids (Mav202 derived). The bait-expressing and preyexpressing strains were to be of opposite mating types. In order to test the two-hybrid interaction between a given bait and prey, the two strains would be mated and the resulting diploid would be selected for on medium appropriate for detecting the interaction (-trp-leu-his, for example). This set up could be scaled up so that a large number of matings were screened simultaneously. However, this protocol required the use of MaV203 derived strains and we had determined that those strains were suboptimal for two-hybrid screening with this set of baits. Therefore, we had to change the protocol to allow for the use of PJ69A derived strains, for which we did not have mating partners. Instead we designed a protocol to perform high throughput screens of the mitosis library that I created from the collection of prey plasmids. Using this protocol I was able to screen the library with 96 individual baits in the same experiment. I used this protocol to screen the mitosis two-hybrid library with each of the 324 bait-expressing, PJ69A-derivative strains in my collection. The interacting prey plasmids were identified by sequence, retested in directed two-hybrid experiments, and the raw data recorded.
- 5. Draw preliminary protein-interaction map for mitotis in C. elegans: Once all the two-hybrid experiments were complete, the data was pooled and annotated according to what screen it came from, etc. I also compared my data to data from Dr. Vidal's lab where similar matrix experiments were being done. Using software called Osprey I was able to organize all of the different data points and draw a preliminary interaction map. The osprey program is designed to store yeast two-hybrid data specifically for yeast proteins. This means that the most advanced features, which interface with databases like SGD and organize data according to gene ontology terms, etc, can't be used for my data because the software doesn't recognize the names of C. elegans ORFs. Therefore, a more appropriate program should be used (or developed) so that we can achieve this level of organization and perhaps create a database for this and future data sets. The complete data set was included with a large data set from Dr. Vidal's lab which was published in January (see attached, ref 2).

Figure 1: Preliminary Interaction Map for Mitosis in *C. elegans* Raw two-hybrid data was compiled and analyzed for potential biological relevance according to what was known about each interacting partner and common sense. This map represents only the few interactions which are considered to be the most interesting for mitosis at this time.



Key Research Accompishments

- Generated bait and prey constructs for a set of 324 C. elegans genes with potential roles in the regulation of
 mitosis. Created a mitosis-specific library from the collection of prey plasmids for use in yeast two-hybrid screens.
- Generated a yeast strain for each of the bait constructs in the collection for use in yeast two-hybrid screens.
- Performed deep yeast two-hybrid screens of complete cDNA libraries with several of the most interesting bait constructs.
- Completed the yeast two-hybrid matrix experiment, wherein the potential interaction of each of the baits in the collection was tested against each prey.
- Generated a preliminary protein-interaction map for this set of proteins.

Reportable Outcomes

Created a protein-interaction map, see figure 1.

Conclusions/Discussion

Although many interactions among the collection of candidate genes have bee recorded, it has yet to be determined if any of these are biologically relevant, or are important for mitosis in C. elegans. Obviously, many more experiments need to be done to make any solid conclusions about these data. One serious concern is that very few of the interactions observed include genes that are currently known to be involved in late mitotic events, either through RNAi phenotypes or by sequence homology to yeast genes. It's possible that proteins interactions that function in mitosis are not readily observed in the yeast two-hybrid system, which would be unfortunate. One notable exception is the homolog of Mob1, which produced two interactions with novel proteins with interesting RNAi phenotypes. One of these novel proteins has an obvious human homolog, which may provide a bridge for this study to continue in a mammalian system.

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REPORTS

The presence of these two patterns in both humans and mouse suggests their importance in the evolution of mammalian X chromosomes. Our sample of functional retroposed genes in the mammalian genomes is likely at least an order of magnitude smaller than the actual number (10, 11). Notably, our analyses exclude retrocopies maintaining introns, such as partially processed retrogenes (35) or chimeric genes (36), which would implicate even more genes. Finally, other mechanisms of interchromosomal gene movement are also likely influenced by the aforementioned selective forces. Thus, we expect many more genes to be subject to the gene traffic described herein.

To elucidate the age of retrogene movements, we dated the human duplications involving X-linked parents or retrogenes both by comparison to the mouse genome sequence and by sequence divergence analysis (16). Most copies that escape X linkage (12/15) as well as most copies that obtain X linkage (10/13) originated before the human-mouse split (Fig. 2, tables S7 and S8). Duplicates in the mouse genome show the same pattern, consistent with this notion. Thus, both patterns result from ancient evolutionary forces common to eutherian mammals. However, this process appears to be an ongoing characteristic of eutherian X evolution, because 6/28 events have occurred subsequent to the human-mouse split in the human lineage, 6/33 retropositions have occurred within the past ~80 million years in the mouse lineage, and some of these retroduplicate pairs have high sequence similarity (>95%) at synonymous sites. This chromosome-biased gene origination appears to be an important process actively driving the differentiation of the X chromosome in mammals and suggests that this differentiation is still in progress.

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Supporting Online Material

www.sciencemag.org/cgi/content/full/303/5657/537/

Materials and Methods Tables S1 to S9

Fig. S1

References

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A Map of the Interactome Network of the Metazoan C. elegans

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To initiate studies on how protein-protein interaction (or "interactome") networks relate to multicellular functions, we have mapped a large fraction of the Caenorhabditis elegans interactome network. Starting with a subset of metazoan-specific proteins, more than 4000 interactions were identified from high-throughput, yeast two-hybrid (HT=Y2H) screens. Independent coaffinity purification assays experimentally validated the overall quality of this Y2H data set. Together with already described Y2H interactions and interologs predicted in silico, the current version of the Worm Interactome (WI5) map contains ~5500 interactions. Topological and biological features of this interactome network, as well as its integration with phenome and transcriptome data sets, lead to numerous biological hypotheses.

To further understand biological processes, it is important to consider protein functions in the context of complex molecular networks. The study of such networks requires the availability of proteome-wide protein-protein interaction, or "interactome," maps. The yeast Saccharomyces cerevisiae has been used to develop a eukaryotic unicellular interactome map (1-6). Caenorhabditis elegans is an ideal model for studying how protein networks relate to multicellularity. Here we investigate its interactome network with HT-Y2H.

As Y2H baits, we selected a set of 3024 worm predicted proteins that relate directly or indirectly to multicellular functions (7). Gateway-cloned open reading frames (ORFs) were available in the C. elegans ORFeome 1.1 (8) for 1978 of these selected proteins. Of these, 81 autoactivated the Y2H GAL1::HIS3 reporter gene as Gal4 DNA binding domain fusions (DB-X), and 24 others conferred toxicity to yeast cells. The remaining 1873 baits were screened against two different Gal4 activation domain libraries (AD-wrmcDNA and AD-ORFeome1.0), each with distinct, yet complementary, advantages (7).

We maximized the specificity of the Y2H system by applying stringent experimental and bioinformatics criteria (fig. S1). To eliminate interactions that originated from nonspecific promoter activation, we only considered DB-X-AD-Y pairs if they activated at least two out of three different Gal4-responsive promoters. Positives were subsequently retested in fresh yeast cells, and their AD-Y identities were determined with interaction sequence tags (ISTs) obtained by sequencing the corresponding polymerase chain reaction (PCR) products (9). The AD-Y reading frame was verified for each IST to avoid the recovery of out-of-frame peptides. In total, ~16,000 ISTs were obtained.

Having applied those criteria, we subdivided the interactions into three confidence classes (fig. S1): those that were found at least three times independently and for which the AD-Y junction is in frame ("Core-1," 858 interactions); those in frame found fewer than three times and that passed the retest ("Core-2;" 1299 interac-

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tions); and all other Y2H interactions found in our screens ("Non-Core," 1892 interactions). The Core data set (Core-1 and Core-2) contains 2157 high-confidence interactions between 502 DB-X baits and 1039 AD-Y preys. After collapsing 22 interactions that occur in both DB-X-AD-Y and DB-Y-AD-X configurations, a total of 2135 unique interactions are obtained (table S1). The Non-Core data set contains 1892 interactions between 531 DB-X baits and 1395 AD-Y preys. Altogether, Core and Non-Core constitute the "First-Pass" data set, with a total of 4027 distinct interactions. Out of 2783 and 1505 interactions found with AD-wrmcDNA and AD-ORFeome1.0, respectively, 239 interactions were identified with both libraries.

To estimate the coverage of the HT-Y2H data sets, we manually searched the baits screened here for known interactors in WormPD (10). This search gave rise to 108 interactions, referred to as the "literature" data set (table S1). The Core and Non-Core data sets recapitulated eight and two interactions in this benchmark data set, respectively. Thus, our overall rate of coverage for the First-Pass data set is $\sim 10\%$ [(8 + 2)/108)].

To evaluate the accuracy of the HT-Y2H data sets, we reasoned that interactions detected in two different binding assays are unlikely to be experimental false-positives. A representative sample of Y2H interaction pairs from each of these three subsets (33 for Core-1, 62 for Core-2, and 48 for Non-Core) was randomly selected, and tested in a coaffinity purification (co-AP) glutathione S-transferase (GST) pull-down assay (Fig. 1). Bait and prey ORFs were transiently transfected into 293T cells as GST-bait and Myc-prey fusions, respectively. For potential interaction pairs where both proteins were expressed at detectable levels. the co-AP success rates were 14 out of 17 (82%) for Core-1, 17 out of 29 (59%) for Core-2, and 8 out of 23 (35%) for Non-Core (table S2). These data demonstrate that our three data sets contain a large proportion of highly reliable interactions and corroborate their expected relative qualities.

In addition to experimental screens, we also performed in silico searches for potentially conserved interactions, or "interologs," whose orthologous pairs are known to interact in one or more other species (9, 11). Starting from a high-confidence yeast interaction data set (7), reciprocal best-hit BLAST searches (E-value ≤ 10⁻⁶) were performed against the worm predicted proteome. In all, 949 potential worm interologs were identified, constituting the interologs data set (7). In addition, the Y2H interactome maps that have been previously generated for individual biological processes (including vulval development, protein degradation, DNA damage response, and germline formation) (9, 12–14) were pooled to define the "scaffold" data set. The HT-Y2H, literature, interologs, and scaffold data sets were combined into Worm Interactome version 5 (WI5). containing 5534 interactions and connecting 15% of the C. elegans proteome (table S1). WI5 gives rise to a giant network component of 2898 nodes connected by 5460 edges (Fig. 2A). Similar to other biological networks (15), the worm interactome network exhibits smallworld and scale-free properties (Fig. 2B) (7). This data set also allowed us to analyze whether or not evolutionary recent proteins tend to preferentially interact with each other rather than with ancient proteins. We subdivided the nodes of the network into three classes: 748 proteins with a clear ortholog in yeast ("ancient"), 1314 proteins with a clear ortholog in Drosophila, Arabidopsis, or humans but not in yeast ("multicellular"), and 836 proteins with no detectable ortholog outside of C. elegans ("worm") (7). These three groups seem to connect equally well with each other (Fig. 2C), which suggests that new cellular functions rely on a combination of evolutionarily new and ancient elements, consonant with the classic proposal of evolution as a tinkerer that modifies and adds to pre-existing structures to create new ones (16).

Previous studies have related interactome data with genome-wide expression (transcrip-

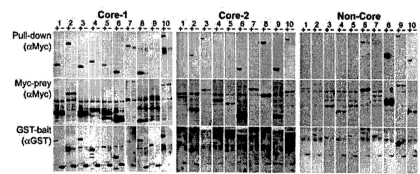


Fig. 1. Coaffinity purification assays. Shown are 10 examples from the Core-1, Core-2, and Non-Core data sets. The top panels show Myc-tagged prey expression after affinity purification on glutathione-Sepharose, demonstrating binding to GST-bait. The middle and bottom panels show expression of Myc-prey and GST-bait, respectively. The lanes alternate between extracts expressing GST-bait proteins (+) and GST alone (-). ORF pairs are identified in table S1 with the lane number corresponding to the order in which they appear in the table.

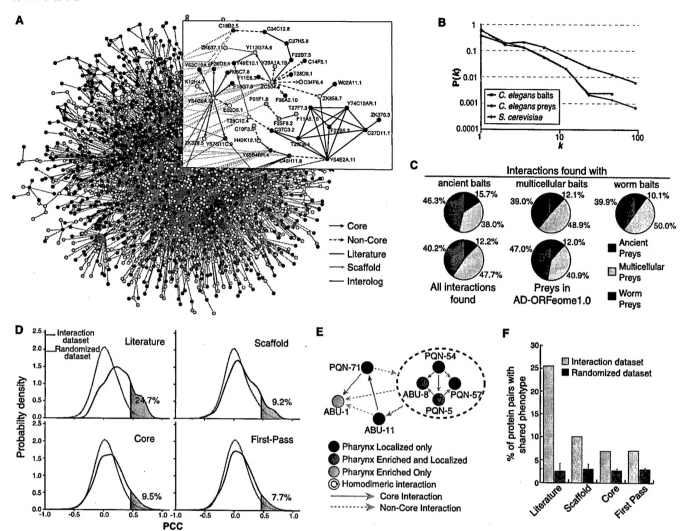


Fig. 2. Analysis of the WI5 network. (A) Nodes (representing proteins) are colored according to their phylogenic class: ancient (red), multicellular (yellow), and worm (blue). Edges represent protein-protein interactions. The inset highlights a small part of the network. (B) The proportion of proteins, P(k), with different numbers of interacting partners, k, is shown for C. elegans proteins used as baits or preys and for S. cerevisiae proteins. (C) The pie charts show the proportion of interacting preys found in Y2H screens that fall into each phylogenic class. Also shown is the distribution of all preys found and all preys searched in the AD-ORFeome 1.0 library.

(D) Overlap with transcriptome (see text) (18), Pearson correlation coefficients (PCCs) were calculated and graphed for each pair of proteins in the interaction data sets and their corresponding randomized data sets. The red area to the right corresponds to interactions that show a significant relationship to expression profiling data (P < 0.05). (E) Interactions between proteins in Topomap mountain 29 (18). The dash-circled proteins belong to the same paralogous family (sharing more than 80% homology) and are thus collapsed into one set of interactions. (F) Proportion of interaction pairs where both genes are embryonic lethal ($P < 10^{-7}$).

tome) and phenotypic profiling (phenome) data in *S. cerevisiae* (17). To investigate to what extent different functional genomic assays should correlate in the context of a multicellular organism, we overlapped WI5 with *C. elegans* transcriptome and phenome data sets.

Based on a C. elegans transcriptome compendium data set (18), we calculated Pearson correlation coefficients (PCCs) for gene pairs involved in Y2H interactions and compared them with randomized data sets (Fig. 2D). About 150 Core interactions (9.5%) corresponded to gene pairs with significantly higher PCCs than expected from random (P < 0.05) (table S3). Thus, those pairs can be considered "more biologically likely" because two completely independent approach-

es point to a functional relationship between the corresponding genes. The remaining pairs are labeled "without additional evidence." Indeed, it is important to note that lack of coexpression does not suggest that the corresponding interactions are irrelevant. Indeed, 75% of literature pairs, defined as biologically relevant, do not correlate with transcriptome data (Fig. 2D).

We also systematically examined Y2H interactions where both proteins belong to common *C. elegans* expression clusters, or "Topomap mountains" (18). As an example, a highly connected subnetwork derived from mountain 29 (Fig. 2E) contains seven proteins (ABU-1, ABU-8, ABU-11, PQN-5, PQN-54, PQN-57, and PQN-71) that share

common domains (DUF139 domain and cysteine-rich repeat). Furthermore, these proteins are all expressed in the pharynx (19-21), which suggests that they may act together in pharynx function or development.

For relatively small-scale S. cerevisiae and C. elegans interactome data sets, physical interactions pointed to genes that share similar phenotypes when knocked out or knocked down (17). To evaluate this idea for the C. elegans interactome, we assembled a collection of phenotypic data based on RNA interference (RNAi) knockdown experiments from WormBase (7, 22), and we calculated the percentage of protein interaction pairs that share embryonic lethal phenotypes for the interaction data sets and their randomized

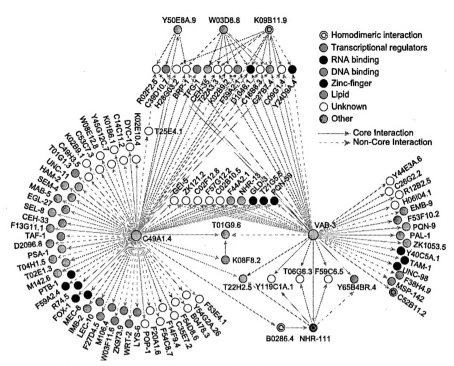


Fig. 3. Graphical representation of a highly interconnected subnetwork around VAB-3 and C49A1.4. Biological functional classes were obtained from WormPD (10).

controls and found a twofold enrichment for the Core and First-Pass data sets (Fig. 2F). Similar correlations were also observed for the maternal sterile phenotype and four groups of postembryonic phenotypes (23). Because protein-protein interactions for which both genes are coexpressed across many conditions and show similar phenotype(s) when knocked down should be considered particularly likely, the global correlations described above illustrate how biological hypotheses can be derived from overlapping interactome, transcriptome, and phenome data sets (table S3).

In S. cerevisiae, two proteins that have many interaction partners in common are more likely to be related biologically (24). We examined the C. elegans interactome network for the presence of highly connected neighborhoods by determining the mutual clustering coefficient between proteins in the network (table S4) (24). As an example, we examined the properties of one of the clusters containing such a highscoring protein pair: VAB-3/C49A1.4 (Fig. 3). VAB-3 and C49A1.4 have strong similarity to the products of the Drosophila genes eyeless (ey) and eyes absent (eya), respectively, but not to each other. EY and EYA are components of a conserved network of transcription factors that regulate eve development (25).

VAB-3 and C49A1.4 are part of a highly interconnected subnetwork in WI5 (Fig. 3) with proteins that are known or suspected to be functionally linked to VAB-3 and C49A1.4, or to their respective orthologs in other organisms. These include (i) EGL-27,

which negatively regulates MAB-5 in hermaphrodites (26) and is linked to MAB-5 through C49A1.4; (ii) WRT-2, an interactor of C49A1.4 with similarity to *Drosophila* Hedgehog, which alleviates repression of eya expression by Cubitus interruptus (27); and (iii) CEH-33 and CEH-35, two of four members of the sine oculis homeobox gene family, which is involved in the same Drosophila regulatory network of transcription factors as ey and eya (28). Finally, eight proteins in this cluster are annotated in WormPD as involved in membrane function, which suggests a functional relationship between the eyeless transcription network and membrane activity.

Together with interologs and previously described interactions, the Y2H data set provides functional hypotheses for thousands of uncharacterized proteins in the C. elegans proteome. Integration with other functional genomic data indicates that the correlation between transcriptome and interactome data, although significant, is lower than what would be expected from observations made in yeast (17). This observation applies to both the Y2H data set described here and wellcharacterized worm interactions from the literature-derived data set (Fig. 2D). This may occur because, unlike unicellular organisms, metazoans are complicated by the fact that biological processes may occur differently in the organism, across various organs, tissues, or single cells.

Our current interactome map also illustrates how a human interactome project would benefit from an ORFeome cloning project using recombinational cloning systems, such as Gateway (δ). Indeed, recombinationally cloned ORFs can be shuffled at will into various expression vectors needed for different types of protein interaction assays, as exemplified by our ability to transfer bait- and prey-encoding ORFs into Myc- and GST-tagged vectors to validate Y2H interactions.

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Supporting Online Material

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Material and Methods

Fig. S1

Table S1 to S5

References

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